## Proposed Decision Memo for Autologous Stem Cell Transplantation (AuSCT) for Amyloidosis (CAG-00050R)

## **Decision Summary**

The Centers for Medicare and Medicaid Services (CMS) proposes the following:

The evidence presented in this decision memorandum is adequate and suggests that when recognized clinical risk factors are employed to select patients for transplantation, high dose melphalan and autologous stem cell transplantation (HDM/AuSCT) can provide a net health benefit for Medicare beneficiaries of any age group with primary AL amyloidosis. HDM/AuSCT is reasonable and necessary for patients with primary AL Amyloidosis who meet the following criteria:

- amyloid deposition in 2 or fewer organs,
- · serum creatinine of 2.0 mg/dL or less, and
- cardiac left ventricular ejection fraction (EF) of 55% or greater.

CMS commends those practitioners who enroll their HDM/AuSCT patients with primary AL amyloidosis in a database (registry). CMS strongly recommends that the sponsors and principal investigators of future HDM/AuSCT trials engage an independent, reputable research center to pool the entire database from each of their respective trials and conduct analyses to identify patient selection, procedure related issues, and other research questions. CMS believes that for optimal patient care, a registry should include criteria that ensure:

- 1. Hospitals and providers are certified as competent in HDM/AuSCT.
- 2. Participating hospitals and providers report data on all patients undergoing HDM/AuSCT.
- 3. Hospitals and providers who do not comply with the data collection requirements are removed from the system.
  - The data set includes elements with the following characteristics:

4.

- Baseline patient characteristics,
- Facility and provider characteristics,
- Extent of disease progression, and
- Long-term patient outcomes.
- 5. Specific hypotheses are addressed.

CMS is requesting public comments on this proposed decision memorandum pursuant to Section 731 of the Medicare Modernization Act. After considering the public comments, we will issue a final decision memorandum.

Back to Top

## **Proposed Decision Memo**

To: Administrative File: CAG-00050R

From:

Steve E. Phurrough, MD, MPA Director, Coverage and Analysis Group

Marcel Salive, MD, MPH Director, Division of Medical and Surgical Services

Samantha Richardson Analyst, Division of Medical and Surgical Services

Susan Harrison Analyst, Division of Medical and Surgical Services

Lori Paserchia, MD Medical Officer, Division of Medical and Surgical Services

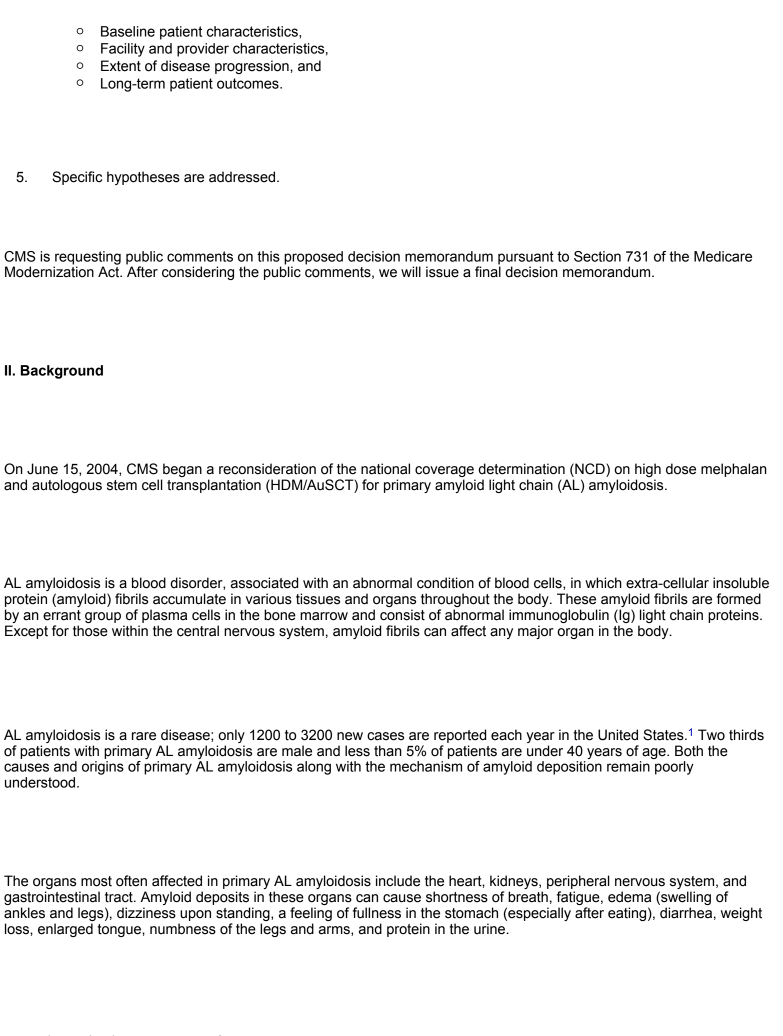
Subject: Proposed Coverage Decision Memorandum for Reconsideration Request for Autologous Stem Cell

Transplantation (AuSCT) for primary amyloid light chain (AL) Amyloidosis

Date: December 15, 2004

Printed on 8/13/2011. Page 2 of 30

l. Prop	oosed Decision
The Ce	enters for Medicare and Medicaid Services (CMS) proposes the following:
factors (HDM/	ridence presented in this decision memorandum is adequate and suggests that when recognized clinical risk are employed to select patients for transplantation, high dose melphalan and autologous stem cell transplantation AuSCT) can provide a net health benefit for Medicare beneficiaries of any age group with primary AL amyloidosis. AuSCT is reasonable and necessary for patients with primary AL Amyloidosis who meet the following criteria:
•	amyloid deposition in 2 or fewer organs, serum creatinine of 2.0 mg/dL or less, and cardiac left ventricular ejection fraction (EF) of 55% or greater.
(registr an inde analys	commends those practitioners who enroll their HDM/AuSCT patients with primary AL amyloidosis in a database ry). CMS strongly recommends that the sponsors and principal investigators of future HDM/AuSCT trials engage ependent, reputable research center to pool the entire database from each of their respective trials and conduct es to identify patient selection, procedure related issues, and other research questions. CMS believes that for I patient care, a registry should include criteria that ensure:
1.	Hospitals and providers are certified as competent in HDM/AuSCT.
2.	Participating hospitals and providers report data on all patients undergoing HDM/AuSCT.
3. 4.	Hospitals and providers who do not comply with the data collection requirements are removed from the system.
	The data set includes elements with the following characteristics:



The clinical course of primary AL amyloidosis is usually associated with rapid disease progression, involvement of multiple organ systems, and short survival periods. Extensive organ system impairment, secondary to amyloid deposits, often results in death. Due to the rapid progression of primary AL amyloidosis, median survival from diagnosis is between one to two years, depending on which organ systems are affected. Patients with cardiac amyloid involvement have an even poorer prognosis with a median survival of less than six months, thus accounting for almost one half of deaths from primary AL amyloidosis.<sup>1</sup>

Because of its similarity to multiple myeloma, another plasma cell dyscrasia where plasma cells produce abnormal protein deposits, the treatment of primary AL amyloidosis has followed the same general path. The early treatment of primary AL amyloidosis largely focused around oral chemotherapy regimens. Patients were treated with standard doses of drugs such as melphalan, prednisone, and/or colchicine. Research suggested that multiple drug regimens can produce better response rates then single drug regimens. However, response rates to standard chemotherapy are quite low. For example, many patients do not live long enough to receive enough cycles of melphalan to actually benefit from treatment.

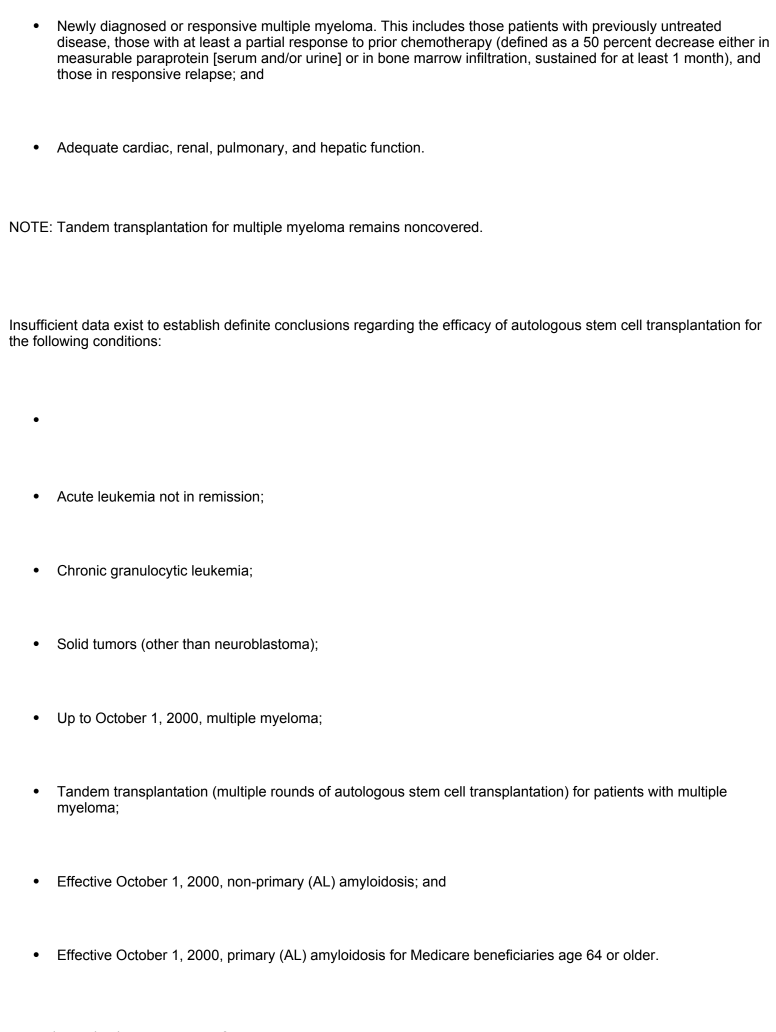
The poor response rates experienced with only chemotherapy prompted the use of HDM/AuSCT. HDM/AuSCT consists of a number of stages. The first stage is called mobilization where the patient is given a granulocyte colony-stimulating factor (G-CSF) or a granulocyte-macrophage colony-stimulating factor (GM-CSF) to stimulate the release of the stem cells from storage sites within the body prior to harvesting via leukapheresis (or bone marrow biopsy). The next stage is called conditioning where the patient is given a high dose of a chemotherapy agent, typically melphalan. In the final stage the harvested stem cells are administered along with supportive medical care.

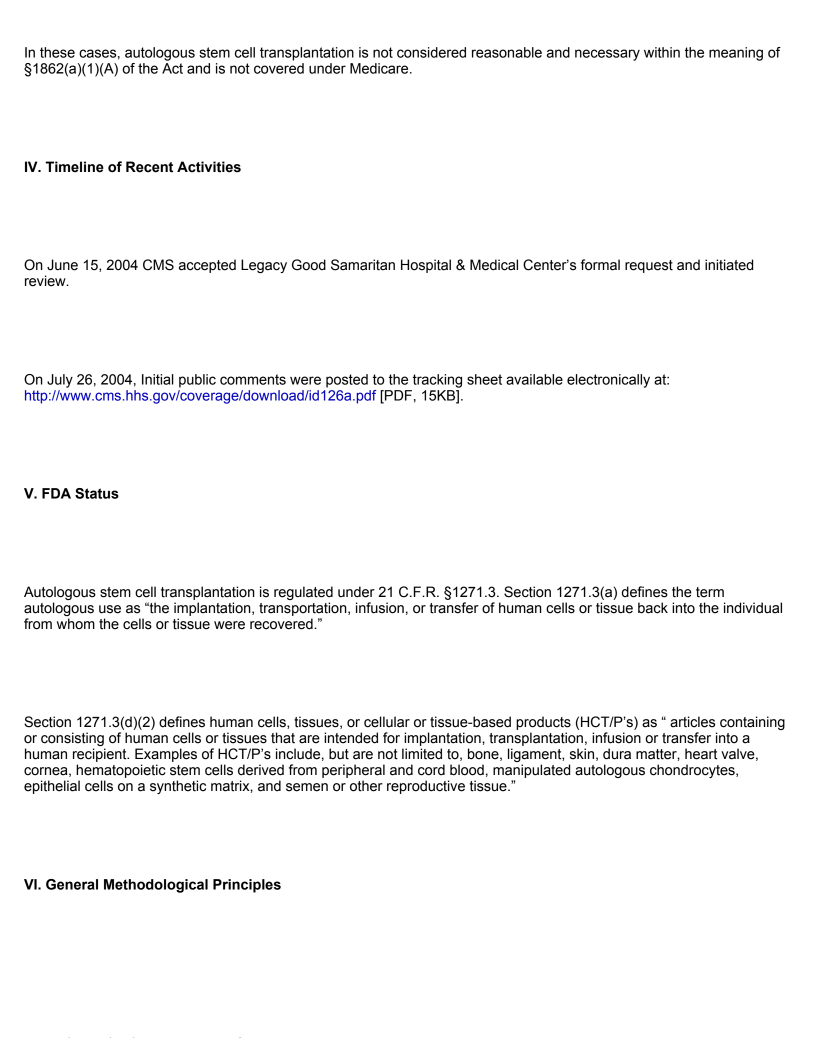
Early on, patients with AL amyloidosis who underwent HDM/AuSCT experienced a reduction in amyloid-related outcomes; however, the transplantation-related mortality was higher compared to patients with multiple myeloma. The presence and severity of amyloid-associated organ dysfunction, which can be extensive in patients with AL amyloidosis and minimal to nonexistent in patients with multiple myeloma, were determined to be the reason for the mortality discrepancy. Subsequent use of HDM/AuSCT in medical practice and studies in research trials permitted the identification of risk factors that guide the selection of patients who are most appropriate to receive HDM/AuSCT. The main factors found to be associated with a significant increased risk of morbidity or mortality include the baseline cardiac and renal status of the patient and the extent of amyloid organ involvement (Comenzo, 2002). Cardiac ejection fraction (EF) was the clinical parameter most commonly used as an inclusion criterion in the clinical trials reviewed for this decision memorandum while the ventricular septal thickness was used less frequently. The serum creatinine level was most commonly used as an inclusion criterion in trials to identify the renal status.

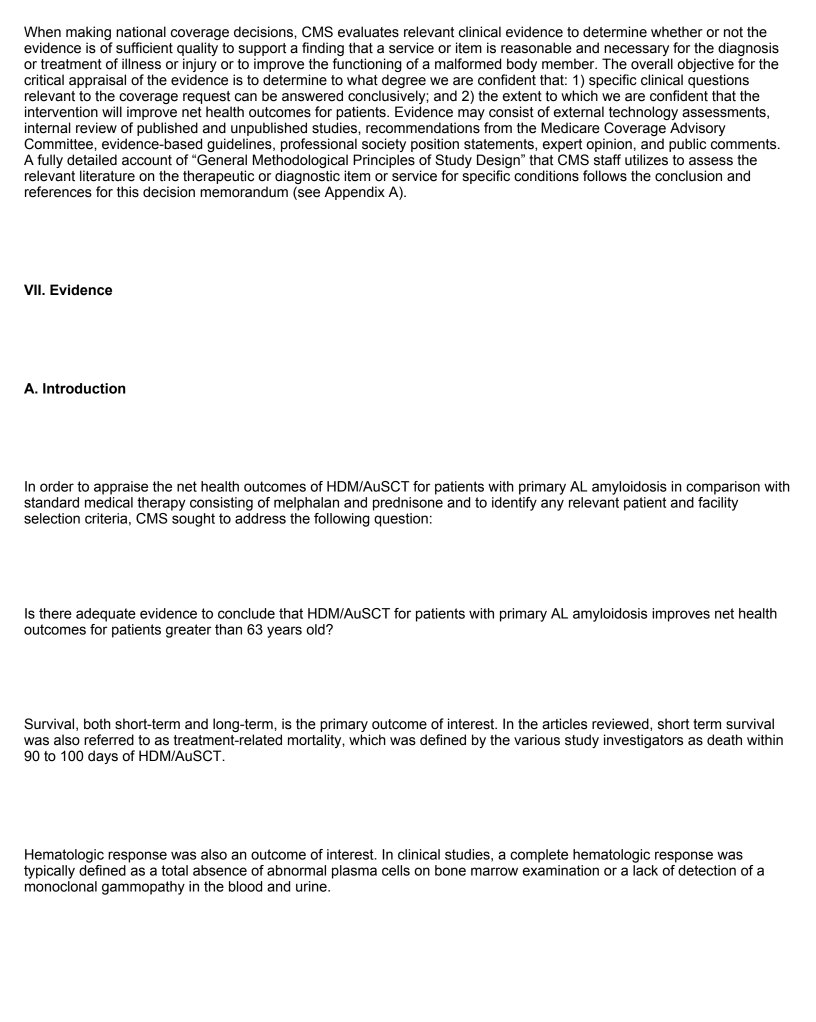
#### III. History of Medicare Coverage

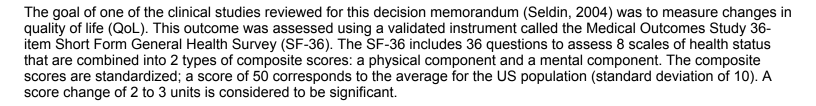
CMS has determined that autologous stem cell transplantation falls within the benefit category of inpatient hospital services under Part A and physicians' services under Part B. See §1812 (a)(1); §1832; §1861(s)(2); §1861(b).

On January 14, 2000, HCFA (now CMS) issued the first decision memorandum (CAG-00050N) which determined that a sufficient body of evidence does not exist to justify a national coverage decision in favor of AuSCT for patients with primary AL amyloidosis. CMS stated that we would reconsider if additional evidence was submitted in the future.
Section 110.8.1 of the National Coverage Determination (NCD) Manual states, in pertinent part, that stem cell transplantation is a process in which stem cells are harvested from either a patient's or donor's bone marrow or peripheral blood for intravenous infusion. The transplant can be used to effect hematopoietic reconstitution following severely myelotoxic doses of chemotherapy and/or radiotherapy used to treat various malignancies.
Autologous stem cell transplantation is considered reasonable and necessary under §1862(a)(1)(A) of the Act for the following conditions and is covered under Medicare for patients with:
•
<ul> <li>Acute leukemia in remission who have a high probability of relapse and who have no human leucocyte antigens (HLA)-matched;</li> </ul>
Resistant non-Hodgkin's or those presenting with poor prognostic features following an initial response;
Recurrent or refractory neuroblastoma; or
Advanced Hodgkin's disease who have failed conventional therapy and have no HLA-matched donor.
Effective October 1, 2000, single AuSCT is only covered for Durie-Salmon Stage II or III patients that fit the following requirement:
•









In Seldin, 2004, the authors noted that QoL instruments exist that are specific for patients undergoing stem cell transplantation. Since patients with primary AL amyloidosis can have a wide range of symptoms due to multi-organ involvement, the SF-36 was selected because it is validated for use in a variety of diseases and patient populations.

#### B. Discussion of evidence reviewed

The evidence reviewed includes summaries of CMS's 2000 AuSCT decision memorandum, CMS's internal technology assessment of new or reconsidered evidence, as well as professional society position statements and expert opinion.

#### 1. Prior CMS Decision Memorandums for AuSCT

In the 2000 decision memorandum "Autologous Stem Cell Transplantation for AL Amyloidosis:" (CAG-00050N), CMS described the etiology of AL amyloidosis and treatments currently available, analyzed relevant clinical literature and delineated reasons for limiting Medicare's current policy of contractor discretion. <sup>2</sup>

In this decision memorandum CMS concluded that a sufficient body of evidence did not exist to justify a NCD in favor of AuSCT for patients with primary AL amyloidosis. The research status appeared to be preliminary and in need of long-term follow-up studies. The majority of AuSCT was performed in highly specialized, typically academic, centers thereby calling in to question the generalizability of the evidence. The clinical studies exhibited a number of deficiencies that increased the risk of bias and confounding, such as small sample sizes and a lack of a randomized control. None of the studies reviewed compared AuSCT to either a control group or other treatment modalities. Selective enrollment of patients resulted in a lack of evidence in patients over 63 years of age and in patients with non-primary AL forms of amyloidosis. Furthermore, the coverage policy in effect at the time for patients younger than 64 years was not revised due to insufficient evidence.

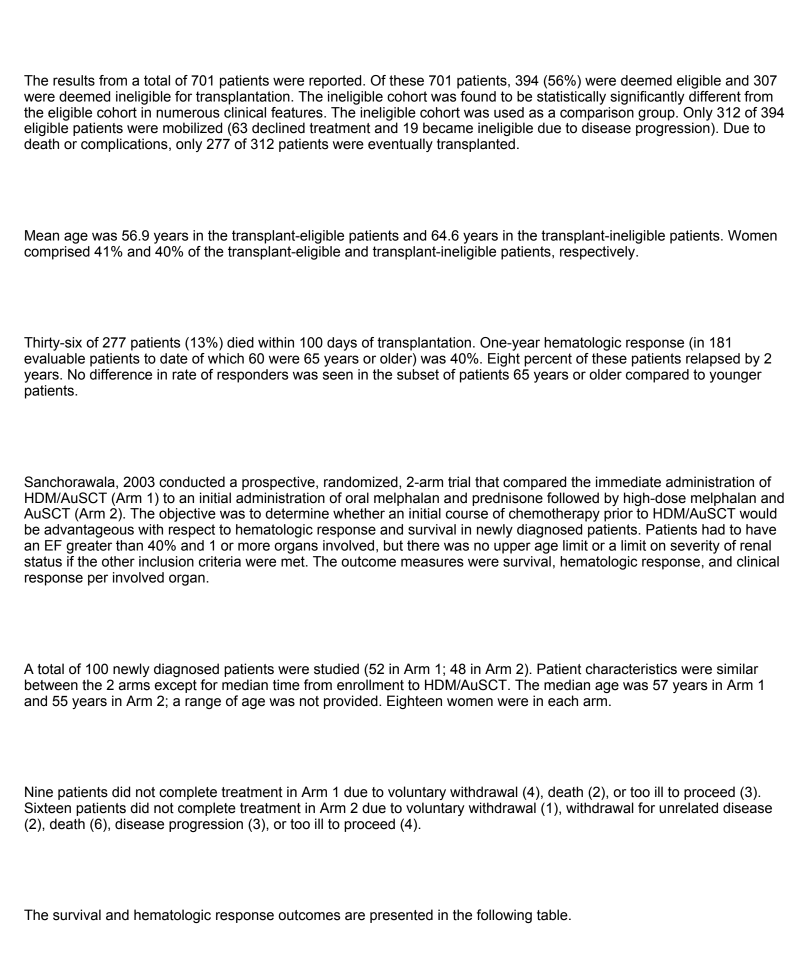
Safety was another concern. A wide range in treatment-related mortality (0, 12%, and 43%) was seen. This emphasized the need for controlled trials and identified proper patient selection as a critical issue to be addressed.

older, and permitted coverage at the discretion of Medicare local contractors for those 63 years old or younger. In addition, CMS decided to not cover AuSCT for Medicare beneficiaries with nonprimary (AL) amyloidosis.
2. External technology assessments
Not applicable.
3. Internal technology assessment
<b>Literature Search</b> CMS performed a search of the literature using the following search terms: peripheral blood, primary amyloidosis, autologous, stem cell transplant. The limitations used were: human, English, Publication date from 1/1/1999 to 11/30/2004. The databases searched were Pub Med, FirstSearch, ProQuest, and EBSCOHost.
Summary of Evidence The requestor submitted 7 published articles and 4 abstracts. The published article for 1 of the abstracts was found and obtained. In addition, the requestor performed and submitted an analysis of the outcome of HDM/AuSCT for patients 65 years of age or older. Sixteen unique abstracts not previously submitted by the requestor were identified based on the following criteria: an abstract was available and the abstract presented the results of a clinical study. Of the 16 abstracts 5 were selected for further review and the full, published article was obtained. Articles subsequently reviewed have either been newly published since CMS's 2000 decision memorandum or are previously published relevant articles now being reconsidered or referenced for the first time.
Scientific articles
In Skinner, 2004, the authors report the pooled results for patients enrolled in 6 distinct protocols. An unblinded, non-

randomized, prospective cohort design was utilized in each study protocol. The inclusion/exclusion criteria and the treatment regimen varied across the protocols. In general, the inclusion criteria permitted patients up to 80 years of age although certain protocols were more age-restricted. Additional inclusion criteria included greater than or equal to 1 major organ involvement, an EF greater than or equal to 40%, and the presence of compensated congestive heart

failure (CHF). Numerous outcomes were measured including survival and complete hematologic response.

Printed on 8/13/2011. Page 11 of 30



Outcomes	Arm 1	Arm 2
Treatment related months: # (0/)		
Treatment-related mortality # (%)		
Pre-stem cell collection	0 (0%)	6 (13%)
Stem cell mobilization/collection	5 (10%)	7 (15%)
Death within 90 days of AuSCT	5 (10%)	4 (8%)
Overall Survival		
1 year	67%	56%
2 year	60%	54%
4 year	51%	50%
5 year	51%	39%
Median Survival (months)	Not reported	37
Complete hematologic response at 1 year	32%	30%

None of the differences were statistically significant.

Seldin, 2004 conducted a prospective, nonrandomized, unblinded, QoL assessment of patients who received HDM/AuSCT. The comparator group consisted of age-matched, transplant-ineligible patients. The purpose of the assessment was to determine if hematologic and clinical responses after HDM/AuSCT are accompanied by an increase in QoL. The outcome was measured using the physical and mental components of the SF-36 form. The hematologic and clinical response outcomes were reported in Skinner, 2004.

Two hundred and fifty-one transplanted patients were compared to 210 age-matched transplant-ineligible patients. The mean age of the transplanted patients was 56±9.5; the mean age for the comparator group was not provided. Patients were mobilized with G-CSF and conditioned with melphalan.

One hundred and four of 251 transplanted patients completed the SF-36 at baseline and at 1 year. There was no apparent difference in clinical characteristics between this group and the 147 patients who did not complete a baseline SF-36. Eighty-four transplanted patients completed the SF-36 at baseline and at 2 years. The presence or absence of differences in clinical characteristics between the 2 groups was not provided. The number of transplant-ineligible patients who completed the SF-36 at baseline and at 1 or 2 years was not provided.

The following table presents the QoL results in the transplanted patients.

Time	Physical Component Score	Mental Component Score
Baseline	34.5	45

Time	Physical Component Score	Mental Component Score
1 year after AuSCT	41	52
2 years after AuSCT	43	51

The physical and mental component scores were not provided for the comparator group. The authors state that quality of life was found to be significantly higher for patients who had a complete hematologic response at 1 year.

A published, peer-reviewed article was not found for 3 of the 4 abstracts submitted by the requestor (Blum, 2001; Lachmann, 2002; Versole, 2003). The evidence table contains a review of these abstracts.

The fourth abstract has been published as a full, peer-reviewed article (Dispenzieri, 2004). In this article the authors report the results of a retrospective case-match-control study of 126 patients (63 cases and 63 controls; 1 case plus 1 control equals a set). Patients who underwent transplantation were matched 1:1 to patients who did not receive transplantation. Matching was based on age, sex, time to presentation, EF, serum creatinine, cardiac septal thickness, nerve involvement, 24-hour urine protein, and serum alkaline phosphatase. The outcomes measured were mortality within 100 days of transplantation and overall survival rate at 1-year, 2-years, and 4-years.

The groups were well matched for age and sex. The median age was 53 years for each group. Four sets of patients were 65 years old or older (range: 66-69 years). The only variables that demonstrated a statistically significant difference between groups at baseline were time from diagnosis to transplantation/treatment (4.4 v. 1.4 months for case v. control, respectively), and EF less than or equal to 50% (6% v. 19% for case v. control, respectively).

Mortality within 100 days of transplantation was 13%. The overall survival rates from the date of transplantation in the case and control groups are presented in the table below.

	Case (n=63)	Control (n=63)
# deaths	16	44
Overall survival rate from transplant date (%)		
1 yr	82*	68
2 yr	81*	53
4 yr	70*	40

Printed on 8/13/2011. Page 14 of 30

Case (n=63)	Control (n=63)

\*P<0.001

Of the 4 case patients greater than or equal to 65 years of age, 1 died at 6.3 months while the remaining 3 case patients are alive after 35, 36, and 38 months. All 4 matched control patients are deceased (death at 6.6, 6.8, 11.6, and 42.6 months).

In Dispenzieri, 2003, a retrospective analysis of the prognostic value of serum cardiac troponin levels is presented. Since this is not a clinical study that focused on the outcome of patients with primary amyloidosis who received HDM/AuSCT, this article was not reviewed.

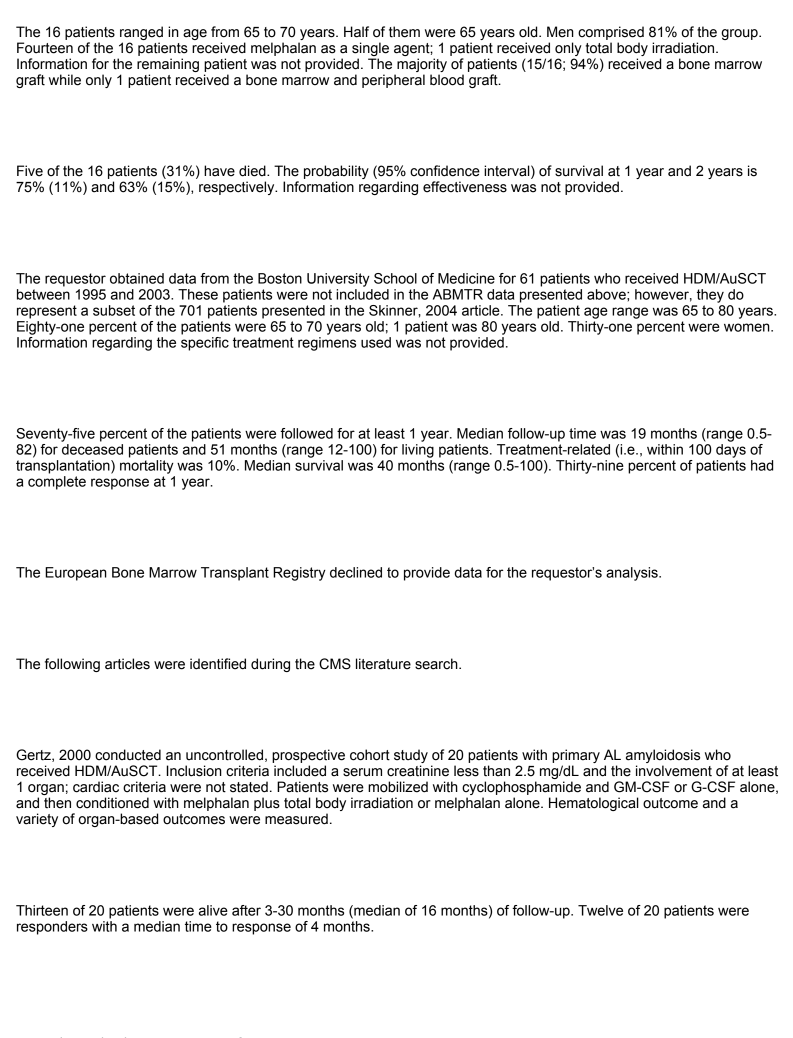
A retrospective case report review was presented in Kumar, 2001. Since this article presents a review of gastrointestinal (GI) bleeding in 4 patients who received HDM/AuSCT, and not a clinical study that focused on determining the clinical outcomes of HDM/AuSCT, it will not be reviewed here.

Similarly, Hayes-Lattin, 2002 is a presentation of 4 case reports focused on the morbidity, especially GI morbidity, associated with HDM/AuSCT. Due to a lack of randomization, blinding, and control found with case reports, robust evidence to support net health outcome decisions is not available; therefore, this article was not considered further.

More details for the 3 articles mentioned above can be found in the evidence table.

CMS asked the requestor to analyze any outcomes data available for patients 65 years old or older who received HDM/AuSCT. The requestor submitted the results of an analysis of data (personal correspondence, Hansen KS and Hansen LK, July 7, 2004) from articles obtained in a Medline search, the Autologous Blood and Marrow Transplant Registry (ABMTR), abstracts from the Tandem Transplant meetings (2002-2004), American Society of Hematology meetings (2001-2003), and the European Bone Marrow Transplant meetings (2001-2003), and from articles posted on the Amyloidosis Support Network website (www.amyloidosis.org).

The ABMTR database contained results for 16 patients who received HDM/AuSCT between 1997 and 2001. The 13 transplant sites were located in North and South America.



Of the 3 patients older than 64 years, 1 patient (70 year old male) died from pneumonia 2 months after transplantation, 1 patient (65 year old male) died from progressive autonomic failure and aspiration 2 months after transplantation, and 1 patient (65 year old male) is living 4 months after transplantation with a complete hematological response.

Dember, 2001 conducted a prospective, uncontrolled cohort trial in 65 patients with renal amyloidosis who were dialysis-independent and at least 18 years old. The objective was to determine the impact of HDM/AuSCT on renal amyloidosis. Inclusion criteria included an EF greater than 40% but the maximal permissible serum creatinine level or the number of involved organs was not stated. Patients were mobilized with G-CSF and conditioned with melphalan, which was dose-adjusted for age, cardiac, renal, pulmonary, and performance status. The outcome measures included 24-hour urinary protein excretion and complete hematologic response.

Median age was 57 years (range 29-77). Women comprised 43% of the patients. Six of 65 patients (9%) died during the peritransplantation period (defined as 100 days after melphalan or during stem cell mobilization or collection). Five of these 6 patients had symptomatic cardiac disease. Fifty of 65 (77%) were alive at 1 year. A comparison of the 1 year survivors versus the non-survivors showed that the survivors were younger (survivor at 1-year median age 56, range 29-71 v. non-survivor at 1-year median age 66, range 40-77, respectively [p=0.024]), had fewer number of involved organ systems, and received a higher melphalan dose. Twenty-one of the 50 1-year survivors had a complete hematologic response. Outcomes for the patients older than 63 years were not separately reported.

The Dispenzieri, 2001 article presents the results of a retrospective case series designed to determine the usefulness of transplant eligibility and other clinical parameters as a prognostic factor. Patients had to have symptomatic organ involvement, be transplant-eligible but not transplanted, and 70 years old or younger. The cardiac septal thickness had to be 15 mm or less, the EF greater than 55%, and the serum creatinine less than or equal to 2.0 mg/dL. Various chemotherapy regimens were administered. Survival was the outcome measure.

A secondary analysis of survival was also performed, which involved a 2:1 case-match-control where the control group was comprised of patients who were transplanted. The match was based on age, sex, and number of involved organs.

Follow-up was available for 96% of the patients. The median time of follow-up was 52 months (range 0.2-186 months). Median survival was 42 months (95% confidence interval, 43-57 months). Survival rates and 95% confidence intervals are presented below.

Time	Case	Control
6 months	83 (75-92)	85 (74-97)
1-year	74 (65-85)	77 (65-91)
2-year	61 (54-68)	68 (53-87)
5-year	36 (30-43)	
10-year	15 (9-24)	

None of these differences were statistically significant.

A number of clinical parameters were found to be predictive of a poor prognosis during both univariate and multivariate analyses: an increasing number of organs involved, worsening performance status, greater than or equal to 10 lb weight loss, and elevation of the serum alkaline phosphatase. Involvement of more than 2 organ systems was associated with worse survival during univariate analysis. Of note, a statistically significant survival difference was not found across the age groups (less than or equal to 50 years, 51-60 years, and 61-70 years).

Gertz, 2002 conducted a prospective, uncontrolled, case series for patients who received HDM/AuSCT between March, 1996 and January, 2001. The use of cardiac or renal status or extent of organ involvement as inclusion criteria was not stated. Sixty-six patients were mobilized with cyclophosphamide and GM-CSF, or with G-CSF alone, and then conditioned with melphalan and total body irradiation, or with melphalan alone. The outcomes measured included complete hematologic response, and various organ-based responses.

The median age was 54 years (range 31-70), and 44% were women. Overall treatment-related mortality was 14%. Thirty -three of 66 (50%) patients had a hematologic response while 32 of 66 (48%) had an organ response. In both univariate and multivariate analyses, serum creatinine and the number of involved organs were associated with mortality.

In Casserly, 2003 the results of a prospective non-randomized, unblinded, concurrent control case series are presented. Patients with amyloidosis-associated end stage renal disease (i.e., dialysis-dependent) who were treated with HDM/AuSCT were included. Patients were excluded for an EF less than 40%, oxygen saturation less than 95% on room air, a performance status greater than or equal to 3, or the presence of refractory CHF or arrhythmia. The extent of organ involvement was not stated as an inclusion/exclusion criterion.

A control group was created that consisted of patients without end stage renal disease who were treated with HDM/AuSCT during the same period. Patients were mobilized with G-CSF alone or with GM-CSF, and conditioned with melphalan. The dose of melphalan was adjusted for age, cardiac, and performance status. The outcome measures were complete hematologic response and survival.

There were 15 cases and 180 control patients. The median age for the cases was 51 years with a range of 40-67; 2 patients were older than 63 years. Females comprised 47% of the population. The demographic profile for the control patients was not provided.

Peritransplant mortality (define as death within 90 days of the start of mobilization) was 13%. The overall hematologic response rate at 1 year was 53% while the hematologic response at 1 year for only the evaluable patients was 73%. Overall median survival was 25 month, and was not statistically significantly different from that in the control group (the survival rate for the control group was not provided).

Printed on 8/13/2011. Page 18 of 30

For the 2 patients who were greater than 63 years, the 67 year old woman had a complete hematologic response and died after 58 months post-transplant due to a hemorrhagic stroke, and the 64 year old woman had a complete hematologic response and is alive after 37 months.
4. MCAC
Not applicable.
5. Evidence-based guidelines
A search of the Web using Google and the search terms "evidence-based guideline" and "stem cell transplantation" yielded no documents. A search of the National Guideline Clearinghouse website was also unsuccessful.
One evidence-based guideline by the National Comprehensive Cancer Network (NCCN) was found via a link from the American Society of Clinical Oncology (ASCO) website (www.asco.org). The 2004 NCCN practice guideline for multiple myeloma briefly notes that insufficient data exist regarding the use of AuSCT for patients with primary AL amyloidosis and "therefore, all patients should be treated in the context of a clinical trial when possible."
A 2004 guideline on the diagnosis and management of amyloidosis by the British Committee for Standards in Hematology notes that HDM/AuSCT may be considered for patients age 70 and under provided the patient has no more than 2 involved organs, does not have a history of amyloidosis-related GI bleeding, and does not have severe cardiomyopathy, advanced renal failure, or is currently on dialysis.
6. Professional Society Position Statements
None found.

#### 7. Expert Opinion

The Comenzo, 2002 article submitted by the requestors is a review of HDM/AuSCT in patients with primary AL amyloidosis. CMS considers this article to be a source of expert opinion.

The authors extensively presented the current status of clinical practice and research, reviewed the AuSCT procedure including peri-transplantation management, and recommended a risk-adapted approach for treating patients. Points highlighted in the article include:

- Median survival of patients seen within 1 month of diagnosis was 13.2 months; less than 5% of all patients with primary AL amyloidosis survive at least 10 years from the time of diagnosis.
- Median survival for patients with CHF was 4 months.
- Patients who undergo HDM/AuSCT typically have a hematologic cancer but no organ dysfunction; patients with primary AL amyloidosis who undergo HDM/AuSCT, on the other hand, typically have multi-organ dysfunction and no cancer.
- Despite previous attempts to define risk-based criteria for patient selection, transplantation-related mortality was still 4-8 times higher in patients with primary AL amyloidosis than in patients with multiple myeloma.
- In addition to the expected risk of chemotherapy-related adverse events during HDM/AuSCT, patients with primary AL amyloidosis also have experienced enhanced toxicity, sometimes fatal, during the mobilization stage of HDM/AuSCT. The cause is unknown. The authors postulate that lower doses of G-CSF or GM-CSF may minimize the risk of toxicity.
- The extent of organ involvement prior to HDM/AuSCT directly influenced the degree of treatment-related mortality.
  - Baseline serum creatinine was a predictor for adverse chemotherapy-related survival and for the transplantation-associated development of renal failure.
  - Based on direct experience, the authors noted a peri-transplantation mortality rate of almost 100% in patients with cardiac amyloid and CHF or with a history of arrhythmia, syncope or recurrent pleural effusion.
- The authors recommended the following risk-adapted approach to selecting and treating patients with primary AL amyloidosis with HDM/AuSCT:

	Good risk (any age; all criteria met)	Intermediate risk (age <71; either criteria)	Poor risk (either criteria)
Extent of organ involvement	1 or 2	1 or 2 (must include cardiac or renal with creatinine clearance <51 mL/min)	<u>&gt;</u> 3
Cardiac involvement	None	Asymptomatic or compensated	advanced
Creatinine clearance	<u>&gt;</u> 51 mL/min		

In a further attempt to adapt treatment to the degree of risk, the authors used the above risk groups as well as the patient's age to guide the dose of melphalan to be administered during the conditioning stage of HDM/AuSCT.

#### 8. Public Comments

CMS received one comment in strong agreement with the information provided by Legacy Good Samaritan Hospital in support of coverage for HDM/AuSCT for patients with primary AL amyloidosis.

#### VIII. CMS Analysis

National coverage determinations (NCDs) are determinations by the Secretary with respect to whether or not a particular item or service is covered nationally under title XVIII of the Social Security Act § 1869(f)(1)(B). In order to be covered by Medicare, an item or service must fall within one or more benefit categories contained within Part A or Part B, and must not be otherwise excluded from coverage. Moreover, with limited exceptions, the expenses incurred for items or services must be "reasonable and necessary for the diagnosis or treatment of illness or injury or to improve the functioning of a malformed body member." § 1862(a) (1) (A).

The quality of the studies of HDM/AuSCT conducted in patients with primary AL amyloidosis since 2000 continues to be less than robust for the Medicare elderly population. Comparative evidence from randomized, controlled trials is not available. While the majority of the studies were prospectively conducted, the majority also were non-randomized. None of the studies were blinded. A control group, when used, was either case-matched (Dispenzieri, 2001; Dispenzieri, 2004) or inappropriate (Sanchorawala, 2003; Seldin, 2004). The requestors, and a number of article authors, state that a randomized, controlled trial will probably never be conducted due to the rarity of the disease and the lack of insurance coverage for patients of Medicare age.

There is some evidence on net health outcomes for patients older than 63 years. Many studies either ultimately did not enroll many patients older than 70 years (Casserly, 2003; Dispenzieri, 2004; Gertz, 2000; Gertz, 2002; Seldin, 2004), intentionally limited enrollment to patients 70 years old or less (Dispenzieri, 2001), or did not specifically note the number of patients older than 63 years (Dember, 2001; Sanchorawala, 2003; Skinner, 2004). The age-related analysis of outcomes performed by the requestor for patients 65 years and older, by our request, provides the most evidence. The majority of patients, however, were 65 to 70 years old; only 19% of the 61 patients were older than 70 years and only 1 was 80 years old.

While most of the studies did not separately report results for patients older than 63 years, the evidence does not suggest worse survival in these patients. The Boston University data from 61 patients who were 65 years and older showed a treatment-related mortality rate of 10%, which compares favorably to the 13% treatment-related mortality rate reported for all 701 patients in Skinner, 2004. A similar overall treatment-related mortality rate was reported in Dember, 2001 (9%), Gertz, 2002 (14%), Casserly, 2003 (13%), and Dispenzieri, 2004 (13%). Dispenzieri, 2004 noted that the survival of the 4 case-matched pairs of patients 65 years old or older was slightly better for the transplant patients than for the non-transplant patients. Compared to the wide range of treatment-related mortality noted in the 2000 decision memorandum, this mortality range across studies is narrower.

Skinner, 2004 reported a median survival of 4.9 years for the subset of patients who were 65 years old and older. This finding was not statistically significantly different from the 4.6 year median survival of the younger patients. Both results contrast with the overall median survival for patients treated with standard chemotherapy of 1-1.5 years (Gertz, 2000). Hence, HDM/AuSCT for patients 63 years of age and older appears to provide a longer survival compared to the currently available treatment.

The univariate and multivariate analyses reported by Dispenzieri, 2001 provide supportive evidence since they found no statistically significant survival difference among the age groups ( $\geq$ 50 years, 51-60 years, and 61-70 years). The multivariate analysis reported in Gertz, 2002 had similar results. Serum creatinine and the number of involved organs, but not age, were found to be independently associated with mortality. The sole study to hint at a survival difference based on the age group was Dember, 2001. Here, a comparison of the survivors and non-survivors at 1-year showed that the survivors were statistically significantly younger.

In contrast to age, the evidence does highlight the relationships between extent of organ system involvement and HDM/AuSCT-related mortality, and between baseline renal status and mortality. Results from analyses performed in Dember, 2001, Dispenzieri, 2001, and Gertz, 2002 point to greater than 2 organ involvement with amyloidosis as strongly associated with mortality, and hence a predictor of a poor prognosis. The Gertz, 2002 analysis found that baseline serum creatinine is independently associated with mortality. These findings serve to support the inclusion and exclusion criteria commonly used to select patients for HDM/AuSCT.

The majority of the studies reviewed for this decision memorandum were conducted in the Stem Cell Transplant Program and the Amyloid Treatment and Research Program at the Boston University School of Medicine in Boston, MA or in the Division of Hematology and Internal Medicine at the Mayo Clinic in Rochester, MN. Additional institutions included the Bone Marrow Transplantation and Leukemia Program at the Washington University School of Medicine in St. Louis, MO; the National Amyloidosis Centre at the Royal Free and University College Medical School in London, UK; the Medical College of Wisconsin in Milwaukee, WI; and the Hematology Service of the Department of Medicine at the Memorial Sloan-Kettering Cancer Center in NY, NY.

The preponderance of major academic institutions in the medical literature signifies the diversity and magnitude of resources necessary to appropriately care for patients with primary AL amyloidosis who undergo treatment with HDM/AuSCT. This point is highlighted by the creation in 1996 of the Foundation for the Accreditation of Cellular Therapy (FACT; www.factwebsite.org) to establish and maintain standards for the safe collection, processing, and administration of hematopoietic cells. It is unlikely that the evidence and results reviewed in this decision memorandum can be readily generalized to facilities with significantly less resources than that typically found at an institution with a hematopoietic transplantation program.

Finally, we desire to ensure that HDM/AuSCT only occurs in those patients who are most likely to benefit and that the procedures are done only by competent providers in facilities with a history of good outcomes and a quality assessment/improvement program to identify providers with poor outcomes and other areas for improvement. As mentioned above, we are concerned that the available evidence does not allow providers to target this therapy to patients who will clearly derive benefit. In order to provide maximum protection to our beneficiaries, CMS will require that reimbursement for HDM/AuSCT occur only if the beneficiary with AL amyloidosis receiving the therapy enrolled in either a FDA approved clinical trial or a qualifying national database (registry).

The submission of surveillance data on patients receiving HDM/AuSCT for AL amyloidosis to a national registry is reasonable and necessary to assure patient safety and protection. Data from the registry will help identify the appropriate patients to receive HDM/AuSCT for AL amyloidosis and help reduce the incidence of inappropriate therapy. These patient protections and safeguards would only be available to the extent that registry data can be made available in some form to providers and practitioners to inform their decisions, monitor performance quality, benchmark and identify best practices. We do not set forth precise standards for data sharing practices. But we do require that the collection and distribution of health information be consistent with the *Standards for Privacy of Individually Identifiable Health Information*.<sup>3</sup>

The national registry for HDM/AuSCT for AL amyloidosis must meet several operational criteria for facility certification, assessment and data completeness. The national registry must include criteria to ensure that hospitals and providers are certified as competent in the AuSCT procedure. Participating hospitals and providers must be required to report data on all patients undergoing HDM/AuSCT for AL amyloidosis. Also, hospitals or providers who do not comply with the data collection requirements must be removed from the system. Complete prospective systematic data collection will ensure the registry's ability to achieve the objectives. Data elements in the national registry should address baseline patient characteristics, facility and provider characteristics, extent of disease progression, and long-term patient outcomes. After the appropriate objectives and hypotheses are developed, the minimum data necessary to answer the hypotheses can be identified, and simple processes for data collection and submission developed.

The registry must be designed to address specific hypotheses, some of which may come from the pooled data analysis of the studies reviewed above. Potential registry hypotheses can center on patient safety-related issues such as:

- the use of risk-based criteria for patient selection as suggested in the Comenzo, 2002 article, and
- the greater degree of G-CSF or GM-CSF associated toxicity seen during the mobilization stage of HDM/AuSCT.

#### IX. Proposed Decision

The Centers for Medicare and Medicaid Services (CMS) proposes the following: The evidence presented in this decision memorandum is adequate and suggests that when recognized clinical risk factors are employed to select patients for transplantation, HDM/AuSCT can provide a net health benefit for Medicare beneficiaries of any age group with primary AL amyloidosis. Based upon the above findings, HDM/AuSCT is reasonable and necessary for patients with primary AL Amyloidosis who meet the following criteria: amyloid deposition in 2 or fewer organs, serum creatinine of 2.0 mg/dL or less, and cardiac left ventricular EF of 55% or greater. CMS commends those practitioners who enroll their HDM/AuSCT patients with primary AL amyloidosis in a database (registry). CMS strongly recommends that the sponsors and principal investigators of future HDM/AuSCT trials engage an independent, reputable research center to pool the entire database from each of their respective trials and conduct analyses to identify patient selection, procedure related issues, and other research questions. CMS believes that for optimal patient care, a registry should include criteria that ensure: Hospitals and providers are certified as competent in HDM/AuSCT. 1. 2. Participating hospitals and providers report data on all patients undergoing HDM/AuSCT. 3. Hospitals and providers who do not comply with the data collection requirements are removed from the system. 4. The data set includes elements with the following characteristics: Baseline patient characteristics, Facility and provider characteristics, Extent of disease progression, and

Long-term patient outcomes.

5. Specific hypotheses are addressed.
CMS is requesting public comments on this proposed decision memorandum pursuant to Section 731 of the Medicare Modernization Act. After considering the public comments, we will issue a final decision memorandum.
Appendix A: General Methodological Principles
We divide the assessment of clinical evidence into three stages: 1) the quality of the individual studies; 2) the relevance of findings from individual studies to the Medicare population; and 3) overarching conclusions that can be drawn from the body of the evidence on the direction and magnitude of the intervention's risks and benefits.
The issues presented here represent a broad discussion of the issues we consider when reviewing clinical evidence. However, it should be noted that each coverage determination has unique methodological aspects.
1. Assessing Individual Studies
Methodologists have developed criteria to determine weaknesses and strengths of clinical research. Strength of evidence generally refers to: 1) the scientific validity underlying study findings regarding causal relationships between health care interventions and health outcomes; and 2) the reduction of bias. In general, some of the methodological attributes associated with stronger evidence include those listed below:
<ul> <li>Use of randomization (allocation of patients to either intervention or control group) in order to minimize bias;</li> <li>Use of contemporaneous control groups (rather than historical controls) in order to ensure comparability between the intervention and control groups;</li> <li>Prospective (rather than retrospective) studies to ensure a more thorough and systematical assessment of</li> </ul>

(intervention or control). This is important especially in subjective outcomes, such as pain or quality of life, where enthusiasm and psychological factors may lead to an improved perceived outcome by either the patient or assessor.

Larger sample sizes in studies to help ensure adequate numbers of patients are enrolled to demonstrate both statistically significant as well as clinically significant outcomes that can be extrapolated to the Medicare population. Sample size should be large enough to make chance an unlikely explanation for what was found; Masking (blinding) to ensure patients and investigators do not know to which group patients were assigned

factors related to outcomes;

Regardless of whether the design of a study is a randomized controlled trial, a non-randomized controlled trial, a cohort study or a case-control study, the primary criterion for methodological strength or quality is the extent to which differences between intervention and control groups can be attributed to the intervention studied. This is known as internal validity. Various types of bias can undermine internal validity. These include:

- Different characteristics between patients participating and those theoretically eligible for study but not participating (selection bias);
- Co-interventions or provision of care apart from the intervention under evaluation (confounding);
- Differential assessment of outcome (detection bias);
- Occurrence and reporting of patients who do not complete the study (attrition bias).

In principle, rankings of research design have been based on the ability of each study design category to minimize these biases. A randomized controlled trial minimizes systematic bias (in theory) by selecting a sample of participants from a particular population and allocating them randomly to the intervention and control groups. Thus, randomized controlled studies have been typically assigned the greatest strength, followed by non-randomized clinical trials and controlled observational studies. The following is a representative list of study designs (some of which have alternative names) ranked from most to least methodologically rigorous in their potential ability to minimize systematic bias:

- Randomized controlled trials:
- Non-randomized controlled trials:
- Prospective cohort studies;
- Retrospective case control studies:
- Cross-sectional studies;
- Surveillance studies (e.g., using registries or surveys);
- Consecutive case series;
- Single case reports.

When there are merely associations but not causal relationships between a study's variables and outcomes, it is important not to draw causal inferences. Confounding refers to independent variables that systematically vary with the causal variable. This distorts measurement of the outcome of interest because its effect size is mixed with the effects of other extraneous factors. For observational, and in some cases randomized controlled trials, the method in which confounding factors are handled (either through stratification or appropriate statistical modeling) are of particular concern. For example, in order to interpret and generalize conclusions to our population of Medicare patients, it may be necessary for studies to match or stratify their intervention and control groups by patient age or co-morbidities.

Methodological strength is, therefore, a multidimensional concept that relates to the design, implementation and analysis of a clinical study. In addition, thorough documentation of the conduct of the research, particularly study's selection criteria, rate of attrition and process for data collection, is essential for CMS to adequately assess the evidence.

#### 2. Generalizability of Clinical Evidence to the Medicare Population

The applicability of the results of a study to other populations, settings, treatment regimens and outcomes assessed is known as external validity. Even well-designed and well-conducted trials may not supply the evidence needed if the results of a study are not applicable to the Medicare population. Evidence that provides accurate information about a population or setting not well represented in the Medicare program would be considered but would suffer from limited generalizability.

The extent to which the results of a trial are applicable to other circumstances is often a matter of judgment that depends on specific study characteristics, primarily the patient population studied (age, sex, severity of disease and presence of co-morbidities) and the care setting (primary to tertiary level of care, as well as the experience and specialization of the care provider). Additional relevant variables are treatment regimens (dosage, timing and route of administration), co-interventions or concomitant therapies, and type of outcome and length of follow-up.

The level of care and the experience of the providers in the study are other crucial elements in assessing a study's external validity. Trial participants in an academic medical center may receive more or different attention than is typically available in non-tertiary settings. For example, an investigator's lengthy and detailed explanations of the potential benefits of the intervention and/or the use of new equipment provided to the academic center by the study sponsor may raise doubts about the applicability of study findings to community practice.

Given the evidence available in the research literature, some degree of generalization about an intervention's potential benefits and harms is invariably required in making coverage decisions for the Medicare population. Conditions that assist us in making reasonable generalizations are biologic plausibility, similarities between the populations studied and Medicare patients (age, sex, ethnicity and clinical presentation) and similarities of the intervention studied to those that would be routinely available in community practice.

A study's selected outcomes are an important consideration in generalizing available clinical evidence to Medicare coverage determinations. One of the goals of our determination process is to assess net health outcomes, and we are interested in the results of changed patient management not just altered management. These outcomes include resultant risks and benefits such as increased or decreased morbidity and mortality. In order to make this determination, it is often necessary to evaluate whether the strength of the evidence is adequate to draw conclusions about the direction and magnitude of each individual outcome relevant to the intervention under study. In addition, it is important that an intervention's benefits are clinically significant and durable, rather than marginal or short-lived.

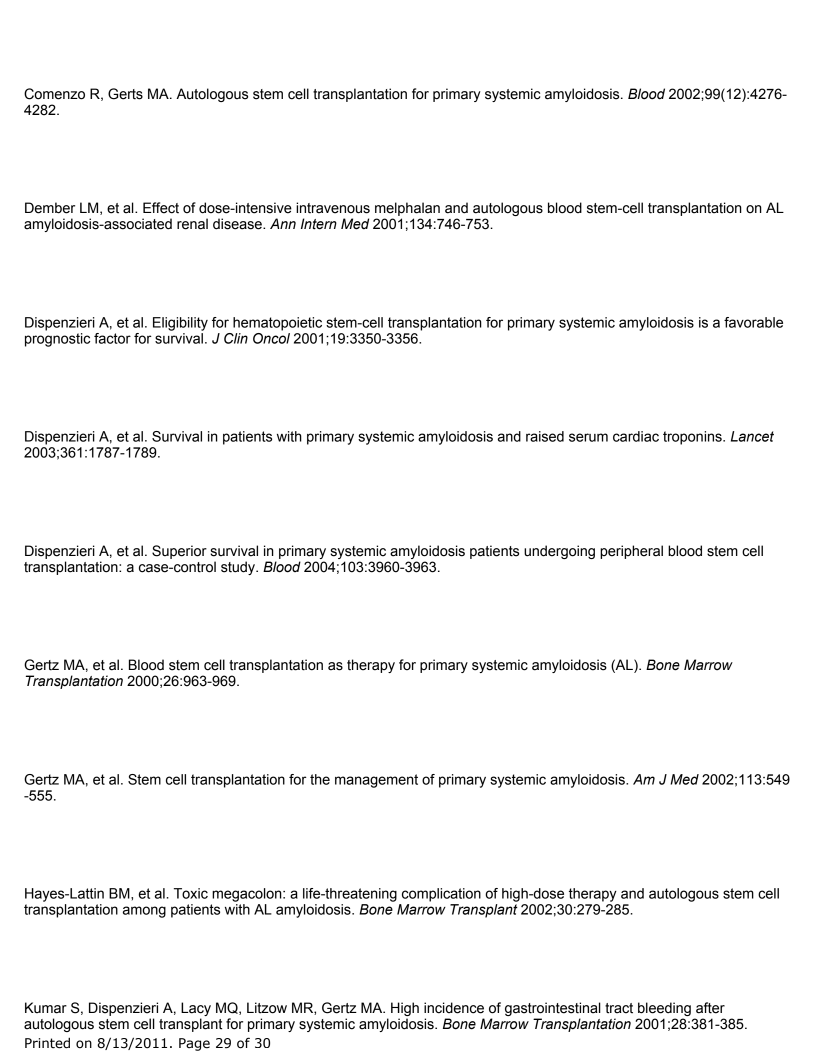
If key health outcomes have not been studied or the direction of clinical effect is inconclusive, we may also evaluate the strength and adequacy of indirect evidence linking intermediate or surrogate outcomes to our outcomes of interest.

# 3. Assessing the Relative Magnitude of Risks and Benefits An intervention is not reasonable and necessary if its risks outweigh its benefits. Among other things, CMS evaluates whether reported benefits translate into improved net health outcomes. The direction, magnitude and consistency of the risks and benefits across studies are important considerations. Based on the analysis of the strength of the evidence. CMS assesses whether an intervention or technology's benefits to Medicare beneficiaries outweigh its harms. Appendix B: Evidence Table [PDF, 219KB] 1 http://amyloidosis.org/treatment/primary.asp#primary <sup>2</sup> http://www.cms.hhs.gov/mcd/viewdecisionmemo.asp?id=9 <sup>3</sup> Privacy Rule – Health Insurance Portability and Accountability Act of 1996. (http://www.os.dhhs.gov/ocr/privacysummary.pdf) Back to Top **Bibliography** Blum W, et al. Primary amyloidosis (AL) patients with significant organ dysfunction treated with conventional chemotherapy followed by single-dose total body irradiation (TBI) and autologous peripheral blood stem cell transplant (PBSC) then IFN-maintenance: tolerance and efficacy. Blood 2001;98(11) abst. 2862:684a.

Casserly LF, et al. High-dose intravenous melphalan with autologous stem cell transplantation in AL amyloidosis-

associated end-stage renal disease. Kidney International 2003;63:1051-1057.

Printed on 8/13/2011. Page 28 of 30



Lachmann HJ, Gillmore JD, Pepys MB, Hawkins PN. Outcome in systemic AL amyloidosis following stem cell transplantation or infusional chemotherapy. <i>Blood</i> 2002:100(11) abstr 788:210a.
Anderson KC, et al. National Comprehensive Cancer Network, Clinical Practice Guidelines in Oncology, Multiple Myeloma, v.1.2005.
Sanchorawala V, et al. High-dose intravenous melphalan and autologous stem cell transplantation as initial therapy or following two cycles of oral chemotherapy for the treatment of AL amyloidosis: results of a prospective randomized trial. Bone Marrow Transplant 2004;33:381-388.
Seldin DC, et al. Improvement in quality of life of patients with al amyloidosis treated with high-dose melphalan and autologous stem cell transplantation. <i>Blood</i> 2004;104:1888-1893.
Skinner M, et al. High-dose and autologous stem-cell transplantation in patients with AL amyloidosis: an 8-year study. Ann Intern Med 2004;140:85-93.
Vesole DH, Perez WS, Reece DE, Akasheh M, Horowitz MM. High dose therapy with autologous hematopoietic stem cell transplantation (HSCT) for patients with primary systemic amyloidosis (AL): results from the autologous blood and marrow transplant registry (ABMTR). <i>Blood</i> 2003:102(11) abst. 402:118a.
Back to Top